A N N U A L   R E P O R T   2 0 1 5

If we don’t care, who will?

www.cfww.org
VISION

All persons living with cystic fibrosis will have access to knowledge and appropriate care.

MISSION STATEMENT

Cystic Fibrosis Worldwide promotes access to knowledge and appropriate care to those people living with cystic fibrosis and among medical, health professionals and governments worldwide.

HISTORY of CYSTIC FIBROSIS WORLDWIDE

Cystic Fibrosis Worldwide (CFW), formed after the merge of the International Association of Cystic Fibrosis Adults and the International Cystic Fibrosis (Mucoviscidosis) Association in 2003, is dedicated to improving quality of life and life expectancy for persons living with cystic fibrosis globally. CFW has 56 member countries with a number of members coming from developing parts of the world. With these new memberships comes an awareness of the desperate situation facing those who have cystic fibrosis, caregivers or medical professionals in developing countries.

CFWW reaches out to these countries by working to help develop effective cystic fibrosis treatment and care. This includes helping to organize much needed medications, development, training and education of health care providers, parents and people with CF and spreading awareness of cystic fibrosis at the government and community levels. We seek to find people with CF who are living in countries where cystic fibrosis is thought to not exist and offer them hope for better tomorrows. It is our objective to find a solution that will bring long-term benefits to the existing persons with cystic fibrosis and to those people with CF who will be born in the future, leaving no people with CF behind. Until a cure is found, we will strive to provide humane situations for those who are currently suffering from cystic fibrosis globally.

What is Cystic Fibrosis?

Cystic fibrosis (CF), also called Mucoviscidosis, is a hereditary disease that affects the entire body, causing progressive disability and early death. Formerly known as cystic fibrosis of the pancreas, this entity has increasingly been termed simply ‘cystic fibrosis.’ Difficulty breathing and insufficient enzyme production in the pancreas are the most common symptoms. Thick mucous production, as well as a low immune system, results in frequent lung infections, which are treated, though not always cured, by oral and intravenous antibiotics, inhalation of mucolytics and hypertonic saline solution, chest physiotherapy and daily exercise. It is vital for CF patients who are pancreatic deficient to use sufficient pancreatic enzymes with every meal and to consume high caloric nutritional foods. A multitude of other symptoms, including sinus infections, poor growth, diarrhea, and potential infertility (mostly in males) result from the effects of CF on other parts of the body.

Approx 1 in 2500 children is born with cystic fibrosis, and it is one of the most common fatal inherited diseases. Individuals with cystic fibrosis can be diagnosed prior to birth by genetic testing through new board screening programs. There is no cure for CF, and most individuals with cystic fibrosis die young — many in their 20s and 30s from lung failure although with many new treatments being introduced the life expectancy for people with CF are increasing. Ultimately, lung transplantation is often necessary as CF worsens.

While CF has existed for a long time the earliest clear medical descriptions date from the 1930’s. Its clinical characteristics individually resemble those of other diseases such as pneumonia, bronchiectasis, failure to thrive, and celiac disease. Indeed, where these conditions are prevalent CF ca go un-recognized. Moreover, if clinicians believe that CF is absent from their population they will not consider it in a differential diagnosis. A better awareness of and the increasing availability of diagnostic tests- the sweat test and/or DNA tests – frequently leads to the identification of a higher number of affected individuals. In the past it was believed CF only existed in Caucasians, through continued advancements and diagnosis we understand CF is a global disease with many different mutations and symptoms.

In the last two decades, CF has been increasingly diagnosed in Latin America, the Middle East, and populations derived from the Indian subcontinent that have emigrated to Western Europe, thus implying the presence of CF in significant numbers among the citizens of India and Pakistan who have remained in their homeland. We now see CF as an emerging disease state throughout the Middle East and Arab populations.

CF is a multi-organ disease and children with CF need access to all necessary medications and multidisciplinary team care in CF centers regularly. This multidisciplinary team should consist of a CF Nurse, Pediatric Pulmonologists, Pulmonologists, Gastroenterologist/Nutritionist, Nurse, Physiotherapist, Microbiologist and Psychologist/Social worker who have been trained in the specific needs of patient care. Daily home care that requires both parent and patient participation is also needed to ensure the patients experience a better quality of life and life expectancy.
Dear friends of Cystic Fibrosis Worldwide.

Another year has gone without a cure found. We will continue to strive for this outcome.

2015 was another busy year for CFW, was a year where we have ensured that we have finalized all report all reporting responsibilities and I can safely say CFW is now in a strong position to strengthen our fundraising drive to support our current and new programs.

With limited funds during 2015 CFW has co-ordinate a Physiotherapy Workshop in Sarajevo during May 2015, we are grateful to members of the CFW Educational team Louise Lannefors and Cecilia Rodriguez-Hortal, for the work they did with the workshop This as you will read in Louise and Cecilia’s report was a very successful workshop and I am sure the people from Bosnia have gained from their professional presentation and support. We also must thank Pari and Horst Melh for their donation in providing nebulizer systems etc and Trip Advisor for their assistance in providing funds for travel and accommodation.

November 2015 CFW coordinated the second CF Latin American two (2) day workshop in Costa Rica, with over 70 people attending, from Costa Rica and 5 other Latin American countries. This workshop was lead by CFW Director Aisha Ramos with 2 Spanish speaking physiotherapist and a dietician , who gave presentations to both medical professionals and lay people and was greatly appreciated by all. There is a third Latin American workshop planned for El Salvador in Nov/Dec 2016.

I have established contact with a private philanthropist in the USA who is interested in providing assistance to Latin American countries. He has already provided support in the region through donations of medicines and equipment and is interested in broadening that involvement I am in regular contact with him to find opportunities for him to increase his involvement.

CFW continues to support our members by supporting their campaigns and projects through letters of support and working to help have CF recognized by health ministries/services. As example of our support is a letter of support we provided to support the Polish Cystic Fibrosis Society application to host a European Cystic Fibrosis Society Conference in the future.

The new CFW website is under development and will be launched in 2016 it is hoped that it will be released in time for the 2016 ECFS Conference in Basel, Switzerland.

I would like to take this opportunity to thank my fellow CFW Board Members for their time, wisdom, knowledge and commitment in making this a better place for the CF Global Community. I also thank our Administrator Danette Lamothe, who has looked after our back of office responsibilities in a very professional way.

To our members and our sponsors, the entire CFW Board thank you for your support and we look forward in working with you during 2016.

_Terry Stewart, CFW Board President and CEO_
Meet the Executive Board of Directors

Terry Stewart
Board President and CEO of Cystic Fibrosis Worldwide
Terry is the former Chief Executive Officer and Company Secretary, Cystic Fibrosis Australia, Chairman of Cystic Fibrosis Australian Data Registry Committee and Cystic Fibrosis Clinical Directors Meetings, Executive Position on all Australasian Cystic Fibrosis Conference Committees, Administrator - Australian Cystic Fibrosis Research Trust
With an open mind on expanding and leading CFW into areas in where CF has limited professional approach to improving the lives of people with Cystic Fibrosis.
Terry founded and ran successful businesses in areas of engineering and property development and brings together a broad wealth of business experience.

Mitch Messer
Treasurer of Cystic Fibrosis Worldwide,
Mitch is an experienced executive, who has served in senior leadership roles as a Chief Executive Officer (CEO) and Board member for a number of organizations over many years. He is a past President of CF Worldwide and currently the CEO of Clan WA, an organization that works to enhance resilience, encouraging healthy relationships, effective parenting and community participation in Western Australia. As a consumer advocate, Mitch is passionate in his belief that we can only truly drive improved health outcomes by ensuring health systems have a consumer focused culture. He believes that CF Worldwide plays an important role in helping to spread knowledge about CF and to finding ways to support and empower people facing the daily challenge of living with CF, especially in parts of the world that do not have the benefit of comprehensive CF care.

Aisha Ramos
Board member CFW Worldwide and Latin America Project committee,
Director of the Spanish CF Association, Federacion Española de Fibrosis Quistica and member of CF Europe. Spain

Horst Mehl
Board member CFW Worldwide, Father of a boy who suffers from CF, Jakob (32 years), Engineer in Baumgartenweg electrotechnics, Diplom-Ingenieur (Dipl. Ing.) D 71364 Winnenden and President of the Mukoviszidose e.V., In den Dauen 6, D 53117 Bonn, since 1996

Catherine Archer
Board Member CFW. Has a family member with CF and brings Global Marketing, Branding skill and experience to our international needs.
Education

“CFW works to educate parents, people living with cystic fibrosis and Medical, Health Professionals in the treatment of cystic fibrosis”

CFW Education Program Team:
Dr. Harry Heijerman, Pulmonologist, CFW Medical Advisor
Dr. Anne Munck, Gastroenterologist, ECFS Board of Directors
Dr. Ian Balfour-Lynn, Royal Brompton Hospital, London
International Nutrition Group/CF
Sue Wolf, ING/CF, Dietician at Leeds Regional Adult CF Unit, UK, member of ECFNG and UK Dieticians
Alison Morton, Dietician at Leeds Regional Adult CF Unit, UK, member of ECFNG and UK Dieticians
International Physiotherapy Group for CF
Louise Lannefors, Physiotherapist, IPG-CF and other IPG/CF therapists
Sue Madge, CF Nurse, International Nurses Group/CF
Maggie McIlwaine, Clinical Investigator, CFRI; Clinical Associate Professor, Physical Therapy, Faculty of Medicine, UBC

Cystic Fibrosis Worldwide is a provider of Cystic Fibrosis (CF) educational programs for practicing clinicians, CFW member associations, patients and families globally with a focus on developing countries. Our Education team of specialist clinical experience is both broad and deep with expertise in inpatient and outpatient treatment of CF to include both well-established and emerging topics in addition to education tailored to primary care practitioners, specialists and allied health professionals.

Cystic Fibrosis Worldwide Educational Program, an integral part of the Cystic Fibrosis Worldwide programs, sponsors conferences, online, and print activities. As a result of these educational activities, Cystic Fibrosis Worldwide fosters the continuing professional development of CF health care providers, CFW member associations, families and patients.

The offerings are designed to promote the professional development of physicians and health care practitioners and influence their clinical practice behavior for the purpose of improving health outcomes for Cystic Fibrosis patients. Families and patients who attend CFW workshops learn how to better cope with CF at home, what questions to ask their caregivers. Physiotherapy, Nutrition and updates on current research and developments help patients and families understand what is needed to give patients a better quality and quantity of life. The CFW Educational program also works to build capacity among our member association via governing, fundraising and program development.

2014-2015 Conferences/Workshops

Annual CFW Meeting and member workshop 2015, Brussels Belgium

Every year, CF Worldwide organizes the Annual Meeting of the Members, in Brussels, where a number members were represented at the CFW ABM.

The CFW Members had a discussion on the CFW interest and involvement in Latin America, advising the members that CFW help co-ordinate the first CF Workshop/Conference in November 2014 within Mexico. This was our first but not our last meeting in Latin America. It was also expressed that this did not mean we were pulling out of Europe, but more a case of broadening our understanding of what could be done in other parts of the world, CFW, does stand for Cystic Fibrosis Worldwide, which means we need to oversee the needs of other areas of the world where CF is prevalent.

Cystic Fibrosis Worldwide
Annual Meeting of the Members
Brussels, Belgium June 2015

Cystic Fibrosis Worldwide - Mexican workshop for Latin America
Dec 2014

Bosnia CF Physiotherapy Course
May 2015

Costa Rica,
November 2015 - 2 day Educational Workshop
Capacity Building

“Cystic Fibrosis Worldwide promotes Capacity Building for CF member associations internationally”

An essential part of CFW programs is to build capacity among our developing member associations. We do this by aiding members through targeted projects. These projects include organizational development workshops, assistance through donations, educational conferences and other various support.

Many new CFW organizations are in developing stages and rely on the international CFW network for guidance and support. CFW helps garner financial resources to aid targeted countries in development stages. Through the CFW Annual Meeting of the Members many ideas, concepts and advice is shared. An open forum fosters continued communications and relationships among CF Associations globally. Through lobbying, clinical development and education along with organizational development support, we reach as many patients as possible with long term solutions.

Thank you Aisha Ramos

Three years ago the CFW Board made the commitment to look at other areas outside Europe that we could help, this was certainly not turning our back on Europe, but more a case of broadening our course to help others where we could. So the decision was made to help the small Latin American countries, based on a needs assessment from information that we received from various organization/individuals in the region.

This of course needed a plan, a contact list of people we could call on to help and someone to drive the initiative. That responsibility fell on the professional shoulders of CFW Board member Aisha Ramos. Aisha is Director of the Spanish CF Association and has worked tirelessly for the CF community.

CFW conducted a segment within the South American mini conference within the 2013 European CF Conference in Lisbon. Aisha presented to the 150 plus delegates who attended the mini conference. A survey was sent out and further information was collected, from there we decided in 2014 to hold a CF Workshop in Mexico City in Nov 2014. This would require a Spanish speaking CF physiotherapist and dietician to visit and present in a 2 day workshop for health professionals and CF parents. The workshop, lead by Aisha Ramos, was very successful. CFW funded travel and accommodation cost of the team to fly from Spain to Mexico City to conduct the workshop. The meeting was well attended and it was also streamed to over 400 people within Mexico, which proved to be a great success for those who could not attend.

Aisha has been an important part of CFW since its inception back in 2002, and has given the organisation the knowledge and ability to get things going in the Spanish speaking Latin American countries.

In November 2015 CFW held its second Latin America Education Workshop in Costa Rica, using the same formula, lead by Aisha with the Spanish speaking professionals from Spain was once again a great success.

We thank Aisha for her ongoing dedication to help bring a better outcome for the CF Community of Latin America. We are also very grateful to two Spanish speaking medical professionals who travelled to both Mexico in 2014 and Costa Rica in 2015 and gave wonderful presentations to both medical professionals and CF parents.

Special thanks to Maria Garriga - Nutritionist & Angeles Fernandez - Physiotherapist
MEXICO REPORT

On the occasion of the IX Congress of the Latin American Society of Pneumology Paediatric (SOLANEP) held in Mexico from 13 to November 16, 2014 we had the opportunity to conduct the First Conference of the CFW training in Latin America.

This conference took place in a parallel workshops and Nutrition Therapy for Cystic Fibrosis parents and professionals who had the sponsorship of the CFW trip through the speakers and broadcasting via streaming of the conference.
A nutritionist, Maria Garriga, a physiotherapist, Angeles Fernández and Aisha Ramos were in charge of the contents of the conference. During the first day the nutritionist gave a full vision to the families while the physiotherapist formed to the professionals, the second day was the other way around. Aisha Ramos spoke to both families and professionals about the work of a national association and the paper of the CFW.

The expectations were surpassed, we had 70 attendees between families and professionals related to CF plus approximately 400 people connected via internet per day, that were able to participate and ask questions to the professionals. This shows the hunger for information with families and patients living with CF. Latin American associations know that there is a long way to go and they were very satisfied for this important step and they need that this event can be repeated in different countries of Latin America.

As the words of the families, they recognize that the information is translated into life, to raise awareness the main responsibility for health care is the family and the patient living with cystic fibrosis and the importance of working together with the staff of the CF centers to achieve significant advances in the quality and life expectancy of people living with Cystic Fibrosis.

After this conference was held the II meeting of the Federation of Cystic Fibrosis Latin America where Mexico took over the presidency for the coming years and where Aisha Ramos participated and pledged to cooperate with them in this new phase of the FLAFQ.

Aisha Ramos, CFW Director
Directora, Federación Española de Fibrosis Quística
C/ Duque de Gaeta 56 pta 14, 46022 Valencia
www.fibrosisquistica.org | Tel. 963318200

THANK YOU TO LUPITA G. CAMPOY

CFW would like to thank Lupita Campoy, who is a Director of Association of CF, Mexico for her co-operation and assistance in helping CFW in delivering an Education Workshop in Mexico City, November 2014.

Furthermore CFW conducted a second CF Latin American Education Workshop in Costa Rica November 2016.

Again, Lupita gave great assistance to Aisha Ramos in setting up this successful Education Workshop. Lupita, who was acting President of the new Cystic Fibrosis Latin America, set up a meeting for all Latin American counties who attended the Costa Rica workshop. It was agreed at that meeting to hold the third Cystic Fibrosis Latin American Education Workshop on the 25,26 and 27 Nov 2016 in Cabo, El Salvador.
During the 2013 CFW Annual Meeting held in Lisbon, Portugal, CFW was approached by the President of the Bosnia CF Association. They asked for a donation of nebulizers, flutters and education on Physiotherapy for patients and families.

Through a grant from TripAdvisor and a donation from board member, Horst Mehl, the program began in 2013, but due to a number of timing issues in holding workshop this will now happen in May 2015.

In May 2015, CFW completed the final phase of the project, hosting a 2 day physiotherapy workshop for families and one day for physiotherapist training in Bosnia.

** Bosnia Physiotherapy Workshop Report - May 28, 2015**

Sarajevo, May 23 - 24, 2015

In Bosnia with ~4 million inhabitants 50 individuals with CF are known, but a big dark number is expected. A few are above 18 years old. The country is divided into 10 cantons, each responsible for organizing their own health care which therefore differs a lot from one canton to another. The knowledge of CF varies extensively. The diagnosis is often suggested by parents who see ill individuals in the hospital, but far too late in the disease progress. Once having had the diagnosis a contact with a physician may be offered, but not for all. Most have never seen a physiotherapist. Drugs, including pancreatic enzymes are often not available within the country and inhalation devices haven’t existed until the recent donation from Pari. But to what extent drugs for inhalation are available varies a lot. Information lacks about optimal use of the nebulizer system, including proper cleaning routines.

The situation of CF care in Bosnia is very difficult due to organization of health care within the country. People are suffering. However families and care givers are eager to learn and spread information about modern CF care, to start organizing adapted, regulated CF care for individuals in different severity levels of the disease, but focusing on preventive care.

**Day 1, parents and patients**

About 35 individuals of whom 4 had CF (1 adult, 2 teenagers and 1 three years old) and 2 physiotherapists from the Sarajevo Paediatric Hospital participated. Several cantons were represented. A translator helped during the whole day. Aiming for a good base the workshop included facts about CF, the progress of the pulmonary disease and the negative spiral the increased respiratory work has on the chest and spine if not taken care of early. Questions on varying levels were brought up and made it obvious how different the knowledge level is. Based on big frustrations patients were eager to discuss their specific problem and having an answer to what is the best treatment for their child – whether what they are doing now is good enough. Much of the prepared material was used, but far from all. Patients attending were instructed and showed how to use the different techniques and the different parts of physiotherapy. A lot of time was spent on questions.

**Day 2, physiotherapists and physicians**

The organizers were not sure about how many participants to expect. Seven arrived; 1 paediatrician, 1 pulmonologist and the chief physiotherapist from the Sarajevo Paediatric Hospital, from another canton 1 paediatrician and 3 physiotherapists in a recently opened pulmonary spa which is now open to CF patients too. A few participants may seem negative, but the questions and discussion became reasonable and fruitful, participants listened to each other. Most of the prepared material was used, focusing on preventive care but also discussing treatment when admitted to hospital. The participants were very engaged. The paediatrician from the spa wanted to be instructed in the use of the different airway clearance techniques. A young CF patient allowed us making it possible to show how to instruct a child.

We were told that participants were satisfied with both days.

Louise Lannefors Copenhagen, Denmark
Cecilia Rodriguez Hortal Stockholm, Sweden
Thank you to the CFW Education Program Team

Louise Lannefors and Cecilia Rodriguez Hortal.

Thank you to Louise Lannefors, who over a number of years has given wonderful service to many physiotherapy workshops for CFW in many different countries.

This has been greatly appreciated by not only by CFW but more importantly by the 1000’s of medical professionals and parents who have participated within these workshops.

Louise is part of the CFW Education Program Team who gives their time and service to the broader CF Communities.

Also a big thank you to Cecilia Rodriguez Hortal, from Stockholm, Sweden, who along with Louise travelled to Bosnia and presented the physiotherapy programs. Cecilia is keen to help out within the CFW Education Program Team and this is a great opportunity for CFW to do further Education workshops in other countries.

**CFW thanks you both for your greatly appreciated service to the CF Community**

Thank you Dr Horst Melh

CFW and the CF Community of Bosnia would like to thank Dr. Horst Melh for his generous donation of the Nebulizers he supplied so the Bosnia Physiotherapy Workshop would go ahead, this was held in Sarajevo, May 2015.

Horst has been a Director of CFW for over 5 years and we great appreciate and respect his wise council in matters dealing with CF in Europe. CFW is honored to have Horst as a Director for his knowledge and his standing within the European CF Community

For many years he has served as President of CF Association Germany, Mukoyiszidose EV, he is currently the Honorary President and Board Member of the CF Association of Germany. He is also a Member of German CF Research Group.
Communication

“Act as a Key Source of international cystic fibrosis information and resources”

CF Worldwide offers an effective platform for the exchange of current information via the Internet in multiple languages for the international CF community at [www.cfww.org](http://www.cfww.org). The Internet is a reliable and convenient information medium for all CFW members/users and medical advisors. The CFW website currently utilizes the very highest standards of web technology, while maintaining a friendly and easy-to-navigate graphic user interface. CFW utilizes feedback from our subscribers and web users to make continued improvements. Along with our active website we also host group pages on Facebook, Twitter and LinkedIn.

New Website

Cystic Fibrosis Worldwide operates a multi-lingual website and online newsletter distributing information about CF globally. We have over 350 translators that help to eliminate language barriers and open lines of communication globally, publications and the latest news and developments in the CF community. The CFW website receives over 100,000 visitors monthly and we look forward to continuing to provide an international platform for the exchange of information.

In 2013, CFW embarked on a project to upgrade our website to make it more user friendly and easier to manage. This was the first major update in many years.

In 2016 we will be updating the site and re-launching it so that it is easier to navigate Websites are always a work in progress and that will continue, as it is a most important element within the CFW communication strategy.

Visit [www.cfww.org](http://www.cfww.org)

Social Media

Social media has become the number one resource for sharing information globally. The CFW Facebook page has 500 likes. Our goal with the CFW Facebook page is to share research and development, member country news, events and personal stories of CF patients in multiple languages. We also share information about the conditions for CF patients in countries around the world.

The CFW Twitter page has almost 1500 followers who share our tweets. We strive to share tweets on a daily basis international news and developments in the CF community.

Plans for 2016

In 2016 we will focus on the value of these programs and review monthly reports on activity of users. We will be focusing on what changes we can make to improve programs or end them if we see they are not a value to the CF international community. This total communication area is now under review to improve this important area for our members and the broader CF community.

Global Voice

“Provide local and international policy and political support to Cystic Fibrosis Associations and people living with cystic fibrosis”

CF Worldwide has a network with partners in the cystic fibrosis global community such as Cystic Fibrosis Worldwide members and international Cystic Fibrosis working groups to bring awareness of cystic fibrosis and the needs of persons with cystic fibrosis at the local and international levels in government, health and public sectors.

CFW lobby’s governments and private sectors of individual countries to ensure support for persons living with cystic fibrosis.
Collaboration

Member Partnerships

CFW has over 60 member countries with a number of members coming from developing parts of the world. With these new memberships comes an awareness of the desperate situation facing those who have cystic fibrosis, caregivers or medical professionals in developing countries. Cystic Fibrosis worldwide members are a continuous support for the many activities carried out by CFW globally. In 2015 some members assisted beyond their annual member dues in order to better facilitate the needs of patients in crises. CFW looks forward to continuing to collaborate with our global network of members in 2016

Our Members

Associate members

Argentina
Australia
Belgium
Canada
Chile
Czech Rep
Denmark
Ecuador
France
Germany
Greece
Hungary
Iceland
Ireland
Latvia
Luxembourg
Macedonia
Netherlands
New Zealand
Norway
Poland
Portugal
Romania
Slovak Republic
Spain
Sweden
Switzerland
Turkey
United Kingdom
United States

Armenia
Austria
Azerbaijan Republic
Bahrain
Brazil
Bulgaria
Costa Rica
Colombia
Dominican Republic
Egypt
El Salvador
Estonia
Finland
Palestine (WB)
Gaza
Italy
India
Jordan
Mexico
Moldova
Lebanon
Lithuania
MATIO
Panama
Paraguay
Qatar
Russia
Saudi Arabia
South Africa
Uruguay
Venezuela
2015 FINANCIAL YEAR SUMMARISED UPDATE

2015 was predominately about stabilising our position and looking at which projects CFW could undertake within our financial restraints.

With the generous funding support from CFW Director Dr Horst Mehl, CFW conducted a Physiotherapy Workshop in Sarajevo, Bosnia in May 2015. This was a great success and within budget.

In December 2014 CFW completed its first Latin American CF Workshop in Mexico City and again within budget.

This was followed up in November 2015 with the second Latin American workshop in Costa Rica. Another success and within budget.

CFW has put money aside to redesign the website. That has been a work in progress for the last 12 months. Again working within a budget.

CFW receives many calls from countries interested in having a CFW sponsored workshop. This is of course subject to our financial position.
2015 Sponsors

Dr Horst Mehl
Germany

- Vertex
- Gilead
- PARI
- TripAdvisor
- Network for Good
- United Way of Massachusetts Bay and Merrimack Valley
“If we don’t care, Who will?”

www.cfww.org